CLAIMS

1. A glycoconjugate comprising a Neisseria meningitidis serogroup B capsular oligosaccharide (MenB OS) derivative in which sialic acid residue N-acetyl groups are replaced with N-acyl groups, wherein said MenB OS derivative is covalently attached to a carrier molecule and has an average degree of polymerization (Dp) of about 10 to about 20.

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- 2. The glycoconjugate of claim 1 wherein the N-acetyl groups are replaced with N-propionyl groups.
- 3. The glycoconjugate of claim 1 wherein the carrier molecule is a bacterial toxoid.
 - 4. The glycoconjugate of claim 3 wherein the bacterial toxoid is tetanus toxoid.
- 5. The glycoconjugate of claim 1 wherein the carrier molecule is a nontoxic mutant bacterial toxoid.
 - 6. The glycoconjugate of claim 5 wherein the mutant bacterial toxoid is CRM_{197} .

- 7. The glycoconjugate of claim 1 wherein the MenB OS derivative has an average Dp of about 12 to about 18.
- 8. A glycoconjugate comprising a Neisseria
 meningitidis serogroup B capsular oligosaccharide (MenB
 OS) derivative in which sialic acid residue N-acetyl
 groups are replaced with N-propionyl groups, wherein said
 MenB OS derivative is covalently attached to a CRM₁₉₇

toxoid protein carrier and has an average Dp of about 12 to about 18.

- 9. The glycoconjugate of claim 1 wherein the MenB OS derivative further comprises a C3-C16 long-chain aliphatic lipid covalently attached thereto.
- 10. The glycoconjugate of claim 8 wherein the MenB OS derivative further comprises a C3-C16 long-chain aliphatic lipid covalently attached thereto.
 - 11. A method for producing a glycoconjugate comprising:
- (a) providing a heterogenous population of

 Neisseria meningitidis serogroup B capsular

 oligosaccharide (MenB OS) derivatives in which sialic
 acid residue N-acetyl groups are replaced with N-acyl
 groups;
- (b) obtaining a substantially homogenous group of MenB OS derivatives from the population of (a) wherein said group of MenB OS derivatives has an average Dp of about 10 to 20;
 - (c) introducing a reactive group at a nonreducing end of the derivatives obtained in step (b) to provide single end-activated MenB OS derivatives; and
 - (d) covalently attaching the end-activated MenB OS derivatives to a carrier molecule to provide a MenB OS glycoconjugate comprising substantially homogenous sized MenB OS moieties.

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12. The method of claim 11 wherein the reactive group introduced in step (c) comprises a reactive aldehyde group.

- 13. The method of claim 11 wherein the sialic acid residue N-acetyl groups of the MenB OS derivatives are replaced with N-propionyl groups.
- 5 14. The method of claim 13 wherein the carrier molecule is a bacterial toxoid.
 - 15. The method of claim 13 wherein the carrier molecule is a nontoxic mutant bacterial toxoid.

- 16. The method of claim 11 wherein the MenB OS derivative has an average Dp of about 12 to about 18.
- 17. The method of claim 11 wherein the MenB OS

 15 derivative further comprises a C3-C16 long-chain

 aliphatic lipid covalently attached thereto.
 - 18. A method for producing a glycoconjugate comprising:
- (a) providing a heterogenous population of Neisseria meningitidis serogroup B capsular oligosaccharide (MenB OS) derivatives in which sialic acid residue N-acetyl groups are replaced with N-propionyl groups;
- of MenB OS derivatives from the population of (a) wherein said MenB OS derivatives have an average Dp of about 12 to 18;
- (c) introducing a reactive group at a nonreducing end of the derivatives obtained in step (b) to provide single end-activated MenB OS derivatives; and
 - (d) covalently attaching the end-activated MenB OS derivatives to a CRM₁₉₇ bacterial toxoid carrier molecule to provide a MenB OS/CRM₁₉₇ toxoid glycoconjugate

comprising substantially homogenous sized MenB OS moieties.

- 19. The method of claim 18 wherein the MenB OS derivative further comprises a C3-C16 long-chain aliphatic lipid covalently attached thereto.
 - 20. A method for producing a glycoconjugate comprising:
- (a) providing a heterogenous population of Neisseria meningitidis serogroup B capsular oligosaccharide (MenB OS) derivatives in which sialic acid residue N-acetyl groups are replaced with N-acyl groups;
- of MenB OS derivatives from the population of (a) wherein said group of MenB OS derivatives has an average Dp of about 10 to 20;
 - (c) introducing a reactive group at a reducing end of the derivatives obtained in step (b) to provide single end-activated MenB OS derivatives; and
 - (d) covalently attaching the end-activated MenB OS derivatives to a carrier molecule to provide a MenB OS glycoconjugate comprising substantially homogenous sized MenB OS moieties.
 - 21. The method of claim 20 wherein the reactive group introduced in step (c) comprises an active ester group.

22. The method of claim 20 wherein the sialic acid residue N-acetyl groups of the MenB OS derivatives

are replaced with N-propionyl groups.

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- 23. The method of claim 22 wherein the carrier molecule is a bacterial toxoid.
- 24. The method of claim 22 wherein the carrier5 molecule is a nontoxic mutant bacterial toxoid.
 - 25. The method of claim 20 wherein the MenB OS derivative has an average Dp of about 12 to about 18.
- 10 26. The method of claim 20 wherein the MenB OS derivative further comprises a C3-C16 long-chain aliphatic lipid covalently attached thereto.
- 27. A method for producing a glycoconjugate 15 comprising:
 - (a) providing a heterogenous population of Neisseria meningitidis serogroup B capsular oligosaccharide (MenB OS) derivatives in which sialic acid residue N-acetyl groups are replaced with N-propionyl groups;
 - (b) obtaining a substantially homogenous group of MenB OS derivatives from the population of (a) wherein said MenB OS derivatives have an average Dp of about 12 to 18;
- 25 (c) introducing a reactive group at a reducing end of the derivatives obtained in step (b) to provide single end-activated MenB OS derivatives; and
 - (d) covalently attaching the end-activated MenB OS derivatives to a CRM₁₉₇ bacterial toxoid carrier molecule to provide a MenB OS/CRM₁₉₇ toxoid glycoconjugate comprising substantially homogenous sized MenB OS moieties.

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- 28. The method of claim 27 wherein the MenB OS derivative further comprises a C3-C16 long-chain aliphatic lipid covalently attached thereto.
- 5 29. A glycoconjugate produced by the method of claim 11.
 - 30. A glycoconjugate produced by the method of claim 18.
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 31. A glycoconjugate produced by the method of claim 20.
- 32. A glycoconjugate produced by the method of claim 27.
 - 33. A vaccine composition comprising the combination of:
- a glycoconjugate formed from a Neisseria

 20 meningitidis serogroup B capsular oligosaccharide (MenB
 OS) derivative in which sialic acid residue N-acetyl
 groups are replaced with N-acyl groups, wherein said MenB
 OS derivative is covalently attached to a carrier
 molecule and has an average degree of polymerization (Dp)
 of about 10 to about 20; and
 - a pharmaceutically acceptable excipient.
- 34. The vaccine composition of claim 33 wherein the N-acetyl groups of the MenB OS derivative are replaced with N-propionyl groups.
 - 35. The vaccine composition of claim 33 wherein the MenB OS derivative has an average Dp of about 12 to about 18.

- 36. The vaccine composition of claim 33 wherein the MenB OS derivative further comprises a C3-C16 long-chain aliphatic lipid covalently attached thereto.
- 37. A vaccine composition comprising the combination of:

a glycoconjugate formed from a Neisseria
meningitidis serogroup B capsular oligosaccharide (MenB
OS) derivative in which sialic acid residue N-acetyl
groups are replaced with N-propionyl groups, wherein said
MenB OS derivative is covalently attached to a CRM₁₉₇
toxoid protein carrier and has an average degree of
polymerization (Dp) of about 12 to about 18; and
a pharmaceutically acceptable excipient.

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- 38. The vaccine composition of claim 37 wherein the MenB OS derivative further comprises a C3-C16 long-chain aliphatic lipid covalently attached thereto.
- 39. The vaccine composition of claim 33 further comprising an adjuvant.
 - 40. The vaccine composition of claim 37 further comprising an adjuvant.

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- 41. A method for preventing Neisseria meningitidis serogroup B and/or E. coli K1 disease in a mammalian subject comprising administering a therapeutically effective amount of the vaccine of claim 33 to said subject.
- 42. A method for preventing Neisseria meningitidis serogroup B and/or E. coli K1 disease in a mammalian subject comprising administering a

therapeutically effective amount of the vaccine of claim 37 to said subject.

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